Amendments to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims:

- 1. (currently amended) A method for identifying biological markers in a set of n biological measurements for each of p observations, wherein n > p and each observation is associated with a clinical endpoint, each biological marker comprising at most k measurements, wherein k < p, said method comprising:
 - a) reducing said set of *n* measurements to a set of *m* candidate measurements; and
 - b) selecting a plurality of at least two biological markers from said set of m candidate measurements;
 - c) determining the accuracy of the said plurality of at least two biological markers in predicting said clinical endpoint; and
 - d) identifying biological markers predictive of wherein values of each biological marker predict-said clinical endpoints.
 - 2. (original) The method of claim 1, wherein said clinical endpoints correspond to clinical classes.
 - 3. (original) The method of claim 1, wherein said clinical endpoints correspond to a continuous response variable.
 - 4. (original) The method of claim 1, wherein n > 10p.
 - 5. (original) The method of claim 1, wherein k < p/5.
 - 6. (original) The method of claim 1, wherein step (a) comprises performing a correlation analysis.
- 7. (original) The method of claim 6, wherein said correlation analysis comprises a correlation-based cluster analysis.

- 8. (original) The method of claim 7, wherein said correlation-based cluster analysis comprises a correlation-based hierarchical cluster analysis.
- 9. (original) The method of claim 6, wherein said correlation analysis is performed in part in dependence on a user-selected correlation threshold.
- 10. (original) The method of claim 6, wherein said correlation analysis is performed in part in dependence on a user-selected value of m.
- 11. (withdrawn) The method of claim 1, wherein step (a) comprises performing a differential significance analysis.
 - 12. (withdrawn) The method of claim 11, wherein said differential significance analysis is performed in part in dependence on a user-selected significance threshold.
- 13. (original) The method of claim 1, wherein said *n* measurements have different sources.
- 14. (original) The method of claim 1, further comprising ranking said selected biological markers.
- 15. (original) The method of claim 14, wherein said biological markers are ranked in dependence on an accuracy of predicting said clinical endpoints.
 - 16. (original) The method of claim 1, wherein said biological markers are selected from all possible subsets of at most k measurements of said set of m measurements.
- 17. (original) The method of claim 16, wherein said biological markers are selected by evaluating each of said possible subsets.

- 18. (original) The method of claim 17, wherein said possible subsets are evaluated in parallel.
- 19. (withdrawn) The method of claim 1, wherein step (b) comprises simulated annealing.
- 20. (original) The method of claim 1, wherein k is a user-selected value.
- 21. (original) The method of claim 1, wherein k is selected in dependence on a desired computation time.
- 22. (original) The method of claim 1, wherein *m* is selected in dependence on a desired computation time.
- 23. (withdrawn) The method of claim 1, further comprising performing a market-basket analysis of said selected biological markers.
- 24. (withdrawn) A method for identifying a biological marker in a set of n biological measurements for each of p observations, wherein n > p and each observation is associated with a clinical endpoint, each biological marker comprising at most k measurements, wherein k < p, said method comprising:
 - a) reducing said set of n measurements to a set of m candidate measurements; and
 - b) using simulated annealing, selecting a biological marker from said set of *m* candidate measurements, wherein values of said biological marker predict said clinical endpoints.
 - 25. (withdrawn) The method of claim 24, wherein n > 10p.
 - 26. (withdrawn) The method of claim 24, wherein k < p/5.

- 27. (withdrawn) The method of claim 24, wherein step (a) comprises performing a correlation analysis.
 - 28. (withdrawn) The method of claim 27, wherein said correlation analysis comprises a correlation-based cluster analysis.
 - 29. (withdrawn) The method of claim 28, wherein said correlation-based cluster analysis comprises a correlation-based hierarchical cluster analysis.
 - 30. (withdrawn) The method of claim 27, wherein said correlation analysis is performed in part in dependence on a user-selected correlation threshold.
 - 31. (withdrawn) The method of claim 27, wherein said correlation analysis is performed in part in dependence on a user-selected value of m.
- 32. (withdrawn) The method of claim 24, wherein step (a) comprises performing a differential significance analysis.
 - 33. (withdrawn) The method of claim 32, wherein said differential significance analysis is performed in part in dependence on a user-selected significance threshold.
- 34. (withdrawn) The method of claim 24, wherein said *n* measurements have different sources.
- 35. (withdrawn) The method of claim 24, wherein k is a user-selected value.
- 36. (withdrawn) The method of claim 24, wherein *k* is selected in dependence on a desired computation time.

- 37. (withdrawn) The method of claim 24, wherein *m* is selected in dependence on a desired computation time.
- 38. (withdrawn) The method of claim 24, further comprising performing a market-basket analysis on said selected biological markers.
- 39. (currently amended) A method for identifying at least one <u>a</u> biological marker in a set of n biological measurements for each of p observations, wherein n > 10p and each observation is associated with a clinical endpoint, each biological marker comprising at most k measurements, wherein k < p, said method comprising:
 - a) reducing said set of n measurements to a set of m candidate measurements; and
 - b) selecting at least one a biological marker from said set of m candidate measurements;
 - c) determining the accuracy of the said at least one biological marker in predicting said clinical endpoint; and
 - d) identifying a biological marker predictive of wherein values of each biological marker predict said clinical endpoints.
- 40. (currently amended) A program storage device accessible by a processor, tangibly embodying a program of instructions executable by said processor to perform method steps for a biological marker identification method, wherein said method identifies biological markers in a set of n biological measurements for each of p observations, wherein n > p and each observation is associated with a clinical endpoint, each biological marker comprising at most k measurements, wherein k < p, said method steps comprising:
 - a) reducing said set of n measurements to a set of m candidate measurements; and
 - b) selecting a plurality of at least two biological markers from said set of m candidate measurements;
 - c) determining the accuracy of the said plurality ofbiological markers in predicting said clinical endpoint; and
 - d) identifying biological markers predictive of wherein values of each biological marker predict said clinical endpoints.

- 41. (withdrawn) A program storage device accessible by a processor, tangibly embodying a program of instructions executable by said processor to perform method steps for a biological marker identification method, wherein said method identifies a biological marker in a set of n biological measurements for each of p observations, wherein n > p and each observation is associated with a clinical endpoint, each biological marker comprising at most k measurements, wherein k < p, said method steps comprising:
 - a) reducing said set of n measurements to a set of m candidate measurements; and
 - b) using simulated annealing, selecting a biological marker from said set of *m* candidate measurements, wherein values of said biological marker predict said clinical endpoints.
- 42. (currently amended) A program storage device accessible by a processor, tangibly embodying a program of instructions executable by said processor to perform method steps for a biological marker identification method, wherein said method identifies \underline{a} at least one biological marker in a set of n biological measurements for each of p observations, wherein n > 10p and each observation is associated with a clinical endpoint, each biological marker comprising at most k measurements, wherein k < p, said method steps comprising:
 - a) reducing said set of n measurements to a set of m candidate measurements; and
 - b) selecting at least one biological marker from said set of m candidate measurements;
 - c) determining the accuracy of the said at least two biological markers in predicting said clinical endpoint; and
 - d) identifying biological markers predictive of wherein values of each biological marker predict said clinical endpoints.
- 43. (previously presented) The method of claim 1, wherein said observations are samples taken at a particular time from a patient belonging to one of a predetermined set of clinical classes.
- 44. (previously presented) The method of claim 43, wherein at least one of said clinical classes is disease condition.

- 45. (previously presented) The method of claim 43, wherein at least one of said clinical classes is response to therapy.
- 46. (new) The method of claim 1, wherein said observations are patients from whom a single or multiple samples are taken.
- 47. (previously presented) The method of claim 1, wherein n is 1000 or more.
- 48. (previously presented) The method of claim 1, wherein n is 5000 or more.
- 49. (previously presented) The method of claim 1, wherein n is 10,000 or more.